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(54) Title: HER2 EXTRACELLULAR DOMAIN

(57) Abstract

An extracellular portion of the HER2 molecule, essentially free of transmembrane and cytoplasmic portions, which is antigenic in animals. Isolated DNA encoding the extracellular portion; an expression vector containing the isolated DNA; and a cell containing the expression vector. A process for producing the extracellular domain. A vaccine containing the extracellular domain.

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HER2 EXTRACELLULAR DOMAIN

BACKGROUND OF THE INVENTION

Field of the Invention

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The present invention is generally directed to the extracellular domain of p185HER2, a receptor-like protein which is encoded by the human homolog of the rat new oncogene.

More specifically, the present invention is directed to a form of the extracellular domain which is essentially free of transmembrane and cytoplasmic domains, to the DNA encoding this form, and to a process for producing this form of the extracellular domain in a host cell.

Description of Background and Relevant Materials

Human epidermal growth factor receptor 2 (HER2, also known as NGL and human c-erbB-2, or ERBB2), is the human homolog of the rat proto-oncogene neu. HER2 encodes a 1,255 amino acid tyrosine kinase receptor-like glycoprotein with homology to the human epidermal growth factor receptor. Although no ligand binding to this probable growth factor receptor has yet been isolated, the HER2 gene product, p185HER2, has the structural and functional properties of subclass I growth factor receptors (Yarden et al., Ann. Rev. Biochem., 57:443-478 (1988); Yarden et al., Biochem., 27:3113-3119 (1988)).

25 The receptor tyrosine kinases all have the same general structural m tif; an extracellular domain that binds ligand, and an intracellular tyrosine kinase d main that is necessary for signal transduction, or in ab rrant

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cases, for transformation. These 2 domains are connected by a single stretch of approximately 20 mostly hydrophobic amino acids, called the transmembrane spanning sequence. This sequence is thought to play a role in transferring the signal generated by ligand binding from the outside of the cell to the inside. It has also been suggested to play a role in the proper positioning of the receptor in the plasma membrane.

Consistent with this general structure, the p185HER2 glycoprotein, which is located on the cell surface, may be divided into three principle portions: an extracellular domain, or ECD (also known as XCD); a transmembrane spanning sequence; and a cytoplasmic, intracellular tyrosine kinase domain. While it is presumed that the extracellular domain is a ligand receptor, as stated above the corresponding ligand has not yet been identified.

The HER2 gene is of particular interest because its amplification has been correlated with certain types of cancer. Amplification of the HER2 gene has been found in human salivary gland and gastric tumor-derived cell lines, gastric and colon adenocarcinomas, and mammary gland adenocarcinomas. Semba et al., Proc. Natl. Acad. Sci. USA, 82:6497-6501 (1985); Yokota et al., Oncogene, 2:283-287 (1988); Zhou et al., Cancer Res., 47:6123-6125 (1987); King et al., Science, 229:974-976 (1985); Kraus et al., EMBO J., 6:605-610 (1987); van de Vijver et al., Mol. Cell. Biol., 7:2019-2023 (1987); Yamam to et al., Nature, 319:230-234 (1986). Gene transfer experiments have shown that

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overexpression of HER2 will transform NIH 3T3 cells and also cause an increase in resistance to the toxic macrophage cytokine tumor necrosis factor. Hudziak et al., "Amplified Expression of the HER2/ERBB2 Oncogene Induces Resistance to Tumor Necrosis Factor Alpha in NIH 3T3 Cells", Proc. Natl. Acad. Sci. USA 85, 5102-5106 (1988).

Because amplification of the HER2 gene results in greatly increased numbers of the p185HER2 protein residing on the surfaces of affected cells, there may interrelationship between increased amounts of p185HER2 extracellular domain on the surfaces of affected cells and the resistance of these cells to treatment. It would therefore be highly desirable to be able to manipulate the p185HER2 extracellular domain in order to investigate several possibilities for the treatment of conditions associated with amplification of the HER2 gene. These therapeutic modes relate not only to the extracellular domain, but also to the putative ligand, which it should be possible to isolate and characterize using the purified p185HER2 extracellular domain.

SUMMARY OF THE INVENTION

The present invention is accordingly directed to an extracellular portion of the HER2 molecule containing at least 9 amino acids, and/or containing an immune epitope, which is essentially free of transmembrane and intracellular portions of the HER2 molecule. The xtracellular portion may be substantially pure, or at least about 99% pure, and may extend to the entire

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extracellular portion of the HER2 molecule. M reover, the extracellular portion may be antigenic in animals, and may be conjugated with a peptide having immunogenic properties; this peptide may contain an immune epitope.

In another embodiment, the present invention is directed to isolated DNA encoding the extracellular portion of the HER2 molecule. This isolated DNA terminates upstream of the DNA portion encoding the transmembrane domain of the HER2 molecule. The termination may occur at least 1 base pair upstream of the portion encoding the transmembrane domain of the HER2 molecule, and preferably occurs about 24 base pairs upstream of this portion.

The isolated DNA of the present invention encodes a sequence of at least 9 amino acids of the extracellular portion, and none of the transmembrane or intracellular portions of the HER2 molecule.

In a further embodiment, the present invention contemplates an expression vector, such as a plasmid or virus, containing the isolated DNA as described above, as well as a cell containing the expression vector. This cell may be eukaryotic or prokaryotic.

The present invention also extends to a process for producing an extracellular portion of the HER2 molecule, which includes the steps of ligating the isolated DNA as described above into an expression vector capable of expressing the isolated DNA in a suitable host; transforming the host with the expression vector;

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culturing the host under conditions suitable for expressi n of the isolated DNA and production of the extracellular portion; and isolating the extracellular portion from the host. The host cell may be a prokaryote, such as a bacterium, or a eukaryote.

In a yet further embodiment, the present invention extends to a vaccine comprising the extracellular portion of the HER2 molecule, which may be combined with suitable adjuvants.

BRIEF DESCRIPTION OF FIGURES

HER2 expression vector and full-length and mutant HER2 proteins. The HER2 expression vector contained eukaryotic transcriptional units for the mouse dihydrofolate reductase (DHFR) cDNA and the bacterial neomycin phosphotransferase (neo) gene, both under SV40 early promoter control. Transcription of the full-length HER2 cDNA was also driven by the early SV40 promoter. full-length HER2 protein contains an extracellular domain with two cysteine-rich clusters (hatched rectangle), separated by the transmembrane-spanning region (filled rectangle) from the intracellular tyrosine kinase domain The mutant protein p185HER2ATM has a (open rectangle). deletion acids. of 28 amino including the transmembrane-spanning region. The truncated p185HERZXCD protein contains all N-terminal sequences up to 8 amino acids before the transmembrane-spanning region.

Fig. 2. Amplification of HER2 and HER2∆TM g nes. Cell lin s transfected with plasmids expressing wild type

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or the ATM mutant HER2 cDNAs were amplified to resistance to 400 nM methotrexate. Cultures were metabolically labeled with [35S]-methionine and proteins immuno-precipitated with the G-H2CT17 antibody. Lane 1: CVN-transfected NIH 3T3 vector control line. Lanes 2 and 3: Parental and amplified HER2-3 line. Lanes 4, 5, and 6, 7: Parent and amplified lines derived from two independent clones, A1 and B2, of the ATM mutant. The arrows indicate the positions expected for proteins of apparent molecular mass of 175 and 185 kDa.

Fig. 3. Autophosphorylation of p185HER2 and p185HER2ATM proteins. Triton X-100 lysates of control, HER2-3400, and ΔTM-expressing cell lines were prepared and immunoprecipitated with the G-H2CT17 antibody. The immune complexes were incubated in 50 ul of HNTG, 5 mM MnC12 with 3 uCi [γ-32P] for 20 min, electrophoresed on a 7.5% polyacrylamide gel, and labeled bands visualized by autoradiography. Lane 1: CVN vector control. Lane 2: HER2-3400 cells expressing full-length HER2. Lanes 3 and 4: Two independent lines expressing p185HER2ATM. The arrows indicate the positions expected for proteins of apparent molecular mass of 66.2, 97, 175, and 185 KDa.

Fig 4. Secretion assay of ΔTM mutants. Cell lines CVN, HER2-3₄₀₀, ΔTM-A1₄₀₀, and ΔTM-B2₄₀₀ were labeled with [³⁵S]-methionine overnight. Triton X-100 cell extracts were prepared and the labeling medium collected. Cells and cell-conditioned media were immunoprecipitated with G-H2CT17 antibody and analyzed on 7.5% SDS-PAGE gels.

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Lanes 1-4 are immunoprecipitations of cell extracts from the various lines, and lanes 5-8 are immunoprecipitations from the corresponding cell-conditioned media. Lanes 1 and 5: CVN vector control. Lanes 2 and 6: HER2-3400 cell lines expressing full-length p185^{HER2}. Lanes 3, 4 and 7, 8: ATM-A1400 and ATM-B2400 cell lines expressing mutant p185^{HER2}. The arrows indicate the positions expected for proteins of apparent molecular mass of 175 and 185 KDa.

Fig 5. Secretion of p185 HER2XCD from 3T3 and CHO cells. NIH 3T3 and CHO cell lines expressing full-length and truncated $p185^{HER2}$ and vector controls were labeled with [35S]-methionine overnight. Cell extracts and cell-conditioned media were immunoprecipitated anti-HER2 monoclonal antibody 3E8 and analyzed on 7.5% SDS-PAGE gels. Lanes 1 and 2: NIH 3T3 control cell line, extract and conditioned medium. Lanes 3 and 4: NIH 3T3 line A1 expressing p185HERZXCD, cells and medium. Lanes 5 and 6: NIH 3T3 line 3400 expressing full-length p185HER2, cells and conditioned medium. Lanes 7 and 8: CHO control line, cell extract and conditioned medium. Lanes 9 and 10: CHO line 2, expressing p185 HERZXCD, cells and conditioned medium. Lanes 11 and 12: CHO line HER2500, expressing full-length p185HER2, cells and conditioned medium. The arrows indicate the molecular mass of the indicated protein bands.

25 Fig 6. Increase in expression of p185HER2XCD with amplification. The CHO-derived cell line HER2XCD-2 was selected for growth in 500 nM and then 3000 nM methotrexate. The parent line, the two amplified

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derivatives, and control vector-transfected cells were labeled with [35S]-methionine. Cell extracts cell-conditioned media were immunoprecipitated with the anti-HER2 monoclonal antibody 3E8 and analyzed on a 7.5% SDS-PAGE gel. Lanes 1 and 2: CVN cell extract and conditioned medium. Lanes 3 and 4: HER2XCD-2, unamplified cells and conditioned medium. Lanes 5 and 6: HER2XCD-2 amplified to resistance to 500 nM methotrexate, cells and conditioned medium. Lanes 7 and 8: HER2XCD-2 amplified to resistance to 3000 nM methotrexate, cells and conditioned medium. For comparative purposes, one-fifth as much sample of the 3000 nm line was loaded compared to the control, 0 and 500 nM lines. The band intensities were quantitated with an LKB2202 laser densitometer. The arrows show the positions of proteins of apparent molecular mass of 88 and 103 KDa.

Fig 7. Biosynthesis of p185HERZXCD. The CHO line HER2XCD23000 was labeled with [35S]-methionine and cell extracts, and cell-conditioned media prepared. Lanes 1 and 2: Cell extract and cell-conditioned medium. Lanes 3 and 4: The same conditioned medium incubated or mock-incubated with endo H. Lanes 5 and 6: Cell extract and conditioned medium from cells treated with tunicamycin. The arrows show the positions expected for proteins of apparent molecular mass of 73, 88, and 103 KDa.

Fig 8. M rphology of NIH 3T3 cells transfected with HER2 and HER2 ATM expression constructs. A and D: Parental and amplified cells fr m NIH 3T3 cells transfected with

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vect r alon . B and E: NIH 3T3 cells expressing p185HER2ATM (line Al), parent and amplified derivative selected for resistance to 400 nM methotrexate. C and F: NIH 3T3 cells expressing wild type p185HER2 (line 3), parent and amplified derivative selected for resistance to 400 nM methotrexate.

Fig 9. Cell surface and cytoplasmic immunofluorescence staining of control, HER2, and HER2ΔTM lines. The top photos are intact cells labeled with anti-HER2 monoclonal antibody. The bottom photos are the same cell lines treated with 0.15% Triton X-100 detergent before addition of antibody. A and D: Control NIH 3T3 cells transfected with vector only. B and E: Cell line HER2 ΔTM-Al₄₀₀, expressing p185HER2ΔTM. C and F: Cell line HER2-3₄₀₀ expressing p185HER2ΔTM.

Fig 10. Fluorescence-activated cell sorter histograms of control, HER2 and HER2ΔTM cells binding anti-p185^{HER2} monoclonal antibody 4D5. Binding by the control antibody, 368, directed against human tissue plasminogen activator, light, broken line. Binding by the anti-HER2 antibody 4D5, dark unbroken line. Panel A: Control NIH 3T3 cells transfected with vector only. Panel B: Cell line HER2-3400, expressing p185^{HER2}. Panel C: Cell line HER2 ΔTMA1400 expressing p185^{ATM}.

Fig 11. Biosynthesis of p185^{HER2} and p185^{HER2} proteins. Cell lines HER2-3₄₀₀ and HER2ATM-A1₄₀₀ were labeled with [³⁵S]-methionine and p185^{HER2} and p185^{HER2} proteins collected by immunoprecipitation and analyzed on a 7.5% SDS-PAGE gel. Lane 1: Vector control. Lane 2: Untreated p185^{HER2} Lanes

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3 and 4: Aliqu ts of the same cell extract treated mock-treated with endo H. Lane 5: Nonglycosylated p185HER2 from cells treated with tunicamycin. Lane 6: Untreated Lanes 7 and 8: Aliquots of the same cell extract treated mock-treated with endo H. Lane 9: p185HERZATH Nonglycosylated from cells treated with tunicamycin. The arrows show the positions of proteins of apparent molecular weight of 175 and 185 kDa.

Purification of the HER2 extracellular Fig. 12. Purified HER2 extracellular domain samples were domain. analyzed using PhastSystem SDS-Gel electrophoresis and silver stained protocols as recommended by Pharmacia. SDS polyacrylamide gel (10-15% gradient) electrophoretic analysis was performed according to Pharmacia protocol File' No. 110. Silver staining was performed according Pharmacia protocol File No. 210. Lane 1 contains molecular weight markers (BRL). Lane 2: Chinese Hamster Ovary Cell 15 X concentrate (1 microliter). Lanes 3 and 4: immunoaffinity purified HER2 extracellular domain (1.6 micrograms and 0.16 microgram, respectively). Lanes 5 and 6: immunoaffinity purified HER2 extracellular domain after DEAE chromatography (0.25 micrograms and 0.083 micrograms, respectively). Lanes 7 and 8: HER2 extracellular domain after formulation in PBS (0.32 micrograms and 0.082 micrograms, respectively).

Fig. 13. The predicted amino acid sequence of the HER2 extracellular d main, with the corresponding nucleic acid sequence. The boxed sequences sh w potential T-cell

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pitopes, using the algorithm developed by Margolit et al.,

J. Immunol. 138:2213-2229(4) (1987).

DETAILED DESCRIPTION

It was initially hypothesized that removal of the transmembrane spanning sequence would yield a protein which would be secreted from the cell. As previously indicated, the transmembrane spanning sequence is principally composed of hydrophobic amino acids, which effectively anchor the protein in the cell membrane. Removal of this sequence would therefore be expected to permit passage of the protein through the membrane.

A first construct was accordingly prepared which deleted exactly in-frame the 22 amino acid transmembrane spanning sequence of HER2, and 3 amino acids on either side (Figure 1). The construct was prepared as follows:

The central EcoR1 fragment containing the transmembrane spanning segment was cloned into the EcoR1 site of the bacteriophage vector M13 mp18 (Yanisch-Perron et al., Gene, 33:103-119 (1985). The noncoding strand was used as template for oligonucleotide-directed mutagenesis. The construct deleted the transmembrane spanning sequence, and an additional 3 amino acids before and after.

Residues 651-678 were deleted by priming double stranded DNA synthesis with a 30 base pair oligonucleotide of sequence 5' CAG AGA GCC AGC CCT CAG CAG AAG ATC CGG 3'. The double stranded DNA was transf rmed into SR101 cells and mutants identified by hybridization to the same oligonucleotide 5' end labeled by polynucleotide kinase and

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 $[\gamma^{-32}P]$ ATP (Amersham, 5000 Ci/mm 1). An Ec R1 fragment containing the deletion was recombined into a plasmid expressing the HER2 cDNA, replacing the wild type sequence.

When expressed in NIH 3T3 cells, this mutant, designated HER2ATM, produced a polypeptide, designated p185HER2ATM, of apparent molecular weight 175 kD (Figure 2, lanes 5 and 7). Production took place at levels comparable to wild type p185HER2 amplified to the same level of resistance to methotrexate (Figure 2, lane 3). The mutant proteins also retained an active tyrosine kinase activity.

In the presence of $[\gamma^{-32}P]$ -ATP, the mutant proteins (Figure 3, lanes 3 and 4) were autophosphorylated to the same extent as unaltered p185 HER (Figure 3, lane 2). Figure 3 also shows autophosphorylated p185HERZATM-related proteins of lower molecular weight than the complete protein. These smaller proteins may represent degradation products and, since they are not observed with p185HER2, could imply a difference in intracellular processing of the mutant form.

> To determine whether the form lacking the transmembrane sequence was secreted, cells were metabolically labeled with 35S-methionine. The culture conditions used herein were as follows: cells were cultured in a 1:1 mixture of Dulbecco's modified Eagle's medium and Ham's nutrient mixture F-12 supplemented with glutamine (2 mM), penicillin (100 units/ml), streptomycin (100 ug/ml), and 10% serum. NIH 3T3-derived cell lines were cultured with calf serum (Hyclone). Chinese Hamster Ovary cells deficient in dihydrofolate reductase (CHO-DHFR) were

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cultured in fetal bovine serum (Gibco) supplement d with glycine (0.13 mM), hypoxanthine (0.11 mM), and thymidine (0.02 mM). (For selection of the transfected plasmid DHFR gene or to amplify introduced plasmids by methotrexate selection, the glycine, hypoxanthine, and thymidine were omitted and extensively dialyzed serum substituted for fetal bovine serum.)

Both cells and cell-conditioned medium were assayed for p185HER2. Figure 4 demonstrates that all p185HER2 remained cell associated (lanes 2, 3, 4), and neither the wild type protein nor the mutant form was secreted (lanes 6, 7, 8).

Thus, contrary to expectations, deletion of the transmembrane spanning sequence was not sufficient to yield a secreted form of $p185^{HER2}$.

The discovery that p185HER^{2aTM} is not secreted suggested that perhaps there are sequences distal to the transmembrane spanning region that prevent passage of p185HER² through the plasma membrane. A second mutant was accordingly made that contained a UAA stop codon 8 amino acids before the beginning of the proposed transmembrane spanning sequence (Figure 1).

The second construct truncated p185^{HER2} 8 amino acids before the start of the transmembrane spanning region at residue 645 by addition of a polypeptide chain-terminating TAA codon. The oligonucleotide 5' AAG GGC TGC CCC GCC GAG TAA TGA TCA CAG AGA GCC AGC CCT 3' was used to prime synthesis of double-stranded DNA from the same template used to construct the ATM mutant. Mutant plaques were

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identified by hybridization to the 5' end-labeled oligonucleotide, and confirmed by checking for the presence of a Bcl 1 site also introduced directly after the ochre codon. The chain-terminated mutant, designated HER2^{XCD}, was then recombined into the HER2 cDNA expression plasmid. The structure of the plasmid and the 2 mutant HER2 derivatives is shown in Figure 1.

Secretion of the resulting form of p185HER2, designated p185HER2KCD, was assayed by first metabolically labeling the cells with 35S-methionine, followed by immunoprecipitation of p185HER2-related proteins from both the cells and cell-conditioned media. In the immunoprecipitation procedure (Hudziak et al., Proc. Natl. Acad. Sci. USA, 84:7159-7163 (1987)), cells were harvested by trypsinization, counted electronically with a Coulter counter, and plated at least 6 hrs. before labeling. plating medium was removed, cells washed with PBS, and the cells re-fed with methionine-free Dulbecco's modified [35S]-methionine (Amersham, 800 Ci/mmol, minimal medium. 29.6 TBq/mmol) was added at 100 uCi/6 cm plate in a volume of 3 ml. Cells were lysed at 4°C with 0.4 ml of HNEG lysis buffer per 6 cm plate. After 10 min, 0.8 ml of lysis dilution buffer (HNEG buffer with 1% bovine serum albumin, 0.1% Triton X-100 detergent) was added to each plate and the extracts were clarified by microcentrifugation for 5 min. Medium to be assayed for secretion of p185HER2 related proteins was collected and clarified by microcentrifugation.

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Antib dies were added to cell extracts or c nditioned medium and allowed to bind at 4°C for 2-4 h. The polyclonal antibody, G-H2CT17(0), recognizing the carboxy-terminal 17 amino acids of p185HERZ, was used for characterization of cell lines expressing the transmembrane-deleted form of The monoclonal antibody 3E8, recognizing an epitope on the extracellular domain (Hudziak et al., Mol. Cell. Bio., 9:1165-1172 (1989)), was used at 8 ug/reaction to immunoprecipitate the truncated form. Seven ug of rabbit anti-mouse IgG was added to immunoprecipitations using this monoclonal to improve its binding to protein A-sepharose. Immune complexes were collected by absorption to protein A-sepharose beads and washed (Hudziak et al., Proc. Natl. Acad. Sci. USA, 85:5102-5106 (1988); Hudziak et al., Proc. Natl. Acad. Sci. USA, 84:7159-7163 (1987)). separated Proteins were on 7.5% sodium sulphate-polyacrylamide gels (SDS-PAGE) and analyzed by autoradiography.

This revealed a form of p185HER2XCD of Mr 88,000 kD that is associated with the cells (Figure 5, lanes 3 and 9); 20 however, the cell-conditioned media from both the NIH 3T3 cells and Chinese hamster ovary-derived lines also contains larger amounts of a protein of Mr 103,000, which is immunoprecipitated by anti-HER2 monoclonal antibody (Figure 5, lanes 4 and 10). Full length p185HER2 was also expressed 25 in both NIH 3T3 and CHO cells (Figure 5), lanes 5 and 11. There is no secretion of native p185HER2 fr m either of these cell types (Figure 5, lanes 6 and 12).

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The larger size f the observed proteins in the cells and cell-conditioned medium (88,000 and 103,000, respectively) compared to the size predicted by the amino acid sequence (71,644) suggested that the truncated form was being glycosylated.

This was confirmed by treating the cells with the antibiotic tunicamycin, which prevents N-linked glycosylation. Treatment with tunicamycin resulted in the appearance of a cell-associated protein of M_r 73,000, which is close to that predicted by the amino acid sequence (Figure 7, lane 5). It also almost completely inhibited secretion of p185HERZXCD into the medium (Figure 7, lane 6). Cell-conditioned medium from tunicamycin treated cells contains only small amounts of the mature 103,000 form, and none of the smaller forms (lane 6). This further suggests that secretion of p185HERZXCD is coupled to glycosylation.

The extent of glycosylation of the secreted form was investigated with the enzyme endoglycanase H (endo H, Boehringer Manheim). This enzyme will hydrolyze asparagine-linked oligosaccharides of the high mannose type. High mannose oligosaccharides are biosynthetic intermediates in the glycosylation process. Final maturation of the carbohydrate side chains involves trimming off some mannose and addition of other sugars such as fucose. Such mature oligosaccharide side chains are resistant to endo H.

To determine if secreted p185HERZKOD is resistant to this enzyme, cell conditioned medium labeled with 35S-m thionine

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was immunoprecipitated. The immuno-precipitates were collected onto protein A sepharose beads and incubated with endo H. Neither mock incubated (lane 3) nor endo H-treated p185 HERZXCD (lane 4) showed any decrease in mobility associated with hydrolysis of the glycosyl side chains, demonstrating that the glycosylation is complete.

Without being bound by any particular theory, these results taken together suggest that the cell-associated form of $p185^{\text{HER2XCD}}$ is an intermediate, and that fully mature glycosylated p185HER2 extracellular domain is being synthesized and secreted. The lack of secretion of the p185HER2aTM protein could be hypothesized to result from the presence of processing information in the transmembrane spanning sequence which is necessary for Golgi transport and targeting of the plasma membrane; however, from these studies it appears instead that transport of tyrosine kinase receptor (or receptor-like) extracellular domain to the cell surface is coupled to proper glycosylation.

Therefore, insertion of the UAA stop codon 8 amino acids before the beginning of the proposed transmembrane spanning sequence yields a fully mature glycosylated p185HER2 extracellular domain which is freely secreted by the cell.

Having succeeded in producing a secreted form of p185HER2, the next stage involved investigating whether the amount of secreted protein could be increased by gene amplification. Using the CHO-d rived cell line, it was found that the amount of extracellular domain could be increased by methotrexate selection. The amount of

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s creted product increased 29-fold in cells selected for resistance to 500 nm methotrexate, and a further 4.4-fold by selection for resistance to 3000 nm methotrexate (Fig. 6).

Thus, a total increase of 128-fold in secreted p185HERZXCD was obtained when this cell line was amplified to resistance to 3000 nm methotrexate, making the production of relatively large quantities of p185HERZXCD possible.

determine whether overexpression of p185 HERZATM results in cell transformation, DNA was introduced in mammalian cells by the CaHPO4 coprecipitation method (Graham et al., Virology, 52:456-467 (1973)). Five ug of plasmid DNA was added to half-confluent plates of cells (6.0 cm) in 1 ml for 4-6 h. The DNA was removed and the cells shocked 15 with 20% (vol/vol) glycerol. After 2 days for phenotypic expression the selective agent geneticin was added at 400 Clones were picked using glass cloning cylinders with petroleum jelly for the bottom seal. The introduced plasmids were amplified by the methotrexate selection procedure (Kaufman et al., J. Mol. Biol., 159:601-621 (1982)).

When the ATM mutant was expressed in NIH 3T3 cells, primary unamplified colonies after selection had the normal flat nontransformed phenotype (Figure 8, compare photo B with vector control alone, photo A). After the expression level was increased by m thotrexate selection, the cells t ok on the refractile, spindle-shaped appearance of transformed cells and also grew piled up in irr gular

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clumps (photo E). This observation is similar to our earlier findings with the unaltered HER2 cDNA (photos C and F, parent and amplified derivatives respectively), and suggests that high levels of expression of the mutant ATM protein were also transforming.

The morphological changes seen at equivalent levels of amplification (400 nm methotrexate) are not as marked for the mutant, implying that the transforming potential of this form of p185 $^{\text{HER2}}$ may be less. At higher levels of resistance to methotrexate, the Δ TM cells become even more transformed in appearance.

The plasmid was also negative in a focus-forming assay whereas the wild type HER2 plasmid was positive, further indicating that the transforming potential of p185HER2ATM protein is lower. Cells expressing high levels also displayed another property of the transformed phenotype, growth in soft agar. Colony formation in soft agar was determined by harvesting each line to be assayed with trypsin, counting the cells (Coulter counter), and plating 80,000 cells per 6-cm dish. The top layer consisted of 4 ml of 0.25% agar (Difco, "purified") over a bottom layer of 5 ml of 0.5% agar. Colonies were counted after 3-4 weeks. Cells from 2 independent clones plated in soft agar gave rise to soft agar colonies with an efficiency comparable to cells expressing the wild type HER2 gene:

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<u>Table I</u> <u>Soft Agar Colony Formation</u>

	<u>Cell Line</u>	# of Soft Agar Colonies
	CVN	0
5	CVN ₄₀₀	. 0
•	HER2-3 ₀	5 +/- 1
	HER2-3 ₄₀₀	208 +/- 27
	ΔTM-A1 ₀	0
	ΔTM-A1 ₄₀₀	205 +/- 62
10	ΔTM-B2 ₀	0
	ΔTM-B2 ₄₀₀	205 +/- 13

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Two control lines were used; NIH 3T3 cells transfected with a plasmid expressing only the neo and DHFR genes, and the same line amplified to resistance to 400 nm methotrexate. The number of soft agar colonies arising was determined for both parental and amplified lines of clones expressing either p185HERZ or p185HERZATM proteins. Each cell line was plated in triplicate and the value averaged.

Therefore, according to the present invention it has been determined that removal of only the transmembrane spanning sequence does not lead to secretion of p185HER2, unless the entire tyrosine kinase domain is also deleted. Removal of this domain results in proper glycosylation and secretion of the extracellular domain.

In order to obtain purified HER2 extracellular domain working material, Chinese Hamster Ovary Cell Harvest Fluid (CFF) containing recombinant HER2 ECD may be first concentrated by ultrafiltration, and then purified by immunoaffinity chromatography using a HER2 specific MAb

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coupled to CNBr activated Sepharose; other suitable immobilization supports may be used. Concentrated CCF is applied to the affinity column after filtration through a 0.2 micron Millipor filter. Purification cycles are performed as necessary until the desired amount of CCF is processed.

During each cycle of purification, the concentrated CCF is applied and the affinity column is washed to baseline with 0.5 M Tris buffer containing 0.15 M NaCl at a pH of approximately 7.5 (TB). HER2 extracellular domain is then eluted from the column with 0.1 M sodium citrate buffer containing 0.5 M NaCl at a pH of approximately 3.5. The affinity column eluant fractions containing HER2 ECD are pooled and neutralized. The immunoaffinity column is reequilibrated between each purification cycle with TB.

In a second step, the affinity column eluant is buffer exchanged into 25 ml of Tris buffer, at a pH of approximately 7.0 (TB2). The HER2 extracellular domain is then applied to a DEAE Sepharose Fast Flow column, and washed with TB2. The HER2 ECD is removed from the column by step or gradient salt elution in TB2 (containing up to 200 mM NaCl).

After DEAE chromatography, purified HER2 ECD fractions are pooled, exchanged into phosphate-buffered saline, and stored at 2-8° C. The resulting material is substantially pure, i. ., about 99% pure (see Fig. 12).

By m ans of the pres nt invention it is accordingly possible to produc a secreted, glycosylated p185HER2

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extracellular d main. This opens several possibilities for further research, as well as a broad range of potential therapeutic applications.

As previously stated, the HER2 gene is of particular interest because its amplification has been correlated with certain types of cancer. In a survey of 189 primary mammary gland adenocarcinomas, it was found that 30% contained amplifications of the HER2 gene. Slamon et al., "Human Breast Cancer: Correlation of Relapse and Survival with Amplification of the HER-2/neu Oncogene," Science 235, 177-182 (1987). Amplification was correlated with a negative prognosis and high probability of relapse.

This suggests that of the 120,000 women diagnosed with breast cancer each year, 36,000 will have HER2 amplification. Approximately half of these women, or about 15,000, may be expected to exhibit greater than 5-fold amplification, corresponding to nearly half of the 40,000 breast cancer-related deaths each year.

It has been demonstrated that a monoclonal antibody against the p185HER2 extracellular specifically inhibits growth of breast tumor-derived cell lines overexpressing the HER2 gene product; prevents HER2transformed NIH 3T3 cells from forming colonies in soft agar; and reduces the resistance to the cytotoxic effect of necrosis factor alpha which accompanies HER2 tumor overexpression. Hudziak <u>et al.</u>, "p185HER2 Monoclonal Antibody has Antiproliferative Effects In Vitro and Sensitizes Human Breast Tumor Cells to Tumor Necrosis

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Factor", Mol. Cell. Biol. 9:1165-1172 (1989). See also, Drebin et al., "Inhibition of Tumor Growth by a Monoclonal Antibody Reactive with an Oncogene-Encoded Tumor Antigen", Proc. Natl. Acad. Sci. USA 83, 9129-9133 (1986) (in vivo treatment with anti-p185 monoclonal antibody asserted to inhibit tumorigenic growth of neu-transformed NIH 3T3 cells implanted in mice).

This effect presents the possibility that conditions characterized by amplification of the HER2 gene may be subject to treatment via Active Specific Immunotherapy. This therapeutic modality contemplates provoking an immune response in a patient by vaccination with an immunogenic form of the extracellular domain. The extracellular domain (or a derivative thereof, as discussed below) may be combined with a local adjuvant which is safe and effective in humans, such as alum, Bacillus calmette-Guerin (BCG), adjuvants derived from BCG cell walls, Detox (Ribiimmunochem), Syntex-1, or Corynebacterium Alternatively, systemic adjuvants, such as Interferon gamma, Interleukin 1, Interleukin 2, or Interleukin 6 may be suitable. An appropriate dose and schedule would be selected to maximize humoral and cell-mediated response.

It may also be possible to enhance an immune response by targeting the immunogen to the immune system, which could lead to more efficient capture of the antigen by antigen presenting cells, or by directing the immunogen so that it is presented by MHC Class 1 mol cules, since th se usually induce a T-cell response.

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In addition to Active Sp cific Immunotherapy, it should be possible to use the purified extracellular domain to isolate and characterize the putative ligand. The HER2 ligand may be used in turn to deliver toxin to tumor cells which are overexpressing HER2, such as by molecular fusion of the ligand with toxin, or by chemical cross-linking. Alternatively, patients overexpressing HER2 may be vaccinated with HER2 ligand conjugated to, or in combination with, a suitable adjuvant.

A patient overexpressing HER2 will also presumably be overexpressing the HER2 ligand. The ligand-HER2 binding interaction, which is likely to contribute to tumor growth, may be inhibited by blocking free ligand in the patient's serum. This blocking can be accomplished by treating the patient with the HER2 extracellular domain, which will proceed to bind free HER2 ligand, thereby preventing the ligand from binding to the HER2 receptor site.

Rather than using the HER2 extracellular domain per se, it may be more desirable to use a derivative which has an increased affinity for the ligand, and/or which has an increased half-life in vivo. Cross-linking on cells is known to improve binding affinity, suggesting that artificial cross-linking can be used to improve the binding ability of the HER2 extracellular domain. The half-life of the extracellular domain in serum can be improved by, for example, fusing the extracellular domain with other molecules present in the serum which are known to have a

long half-life, such as the Fc-p rtion of an immun gl bin molecule.

The present invention has of necessity been discussed herein by reference to certain specific methods and materials. It is to be understood that the discussion of these specific methods and materials in no way constitutes any limitation on the scope of the present invention, which extends to any and all alternative materials and methods suitable for accomplishing the ends of the present invention.

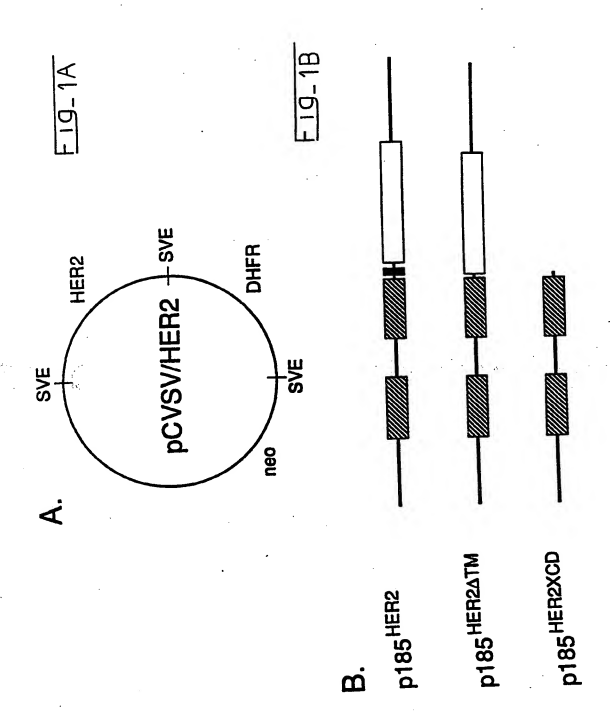
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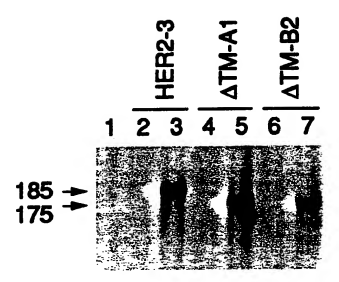
WHAT WE CLAIM IS:

- 1. An extracellular portion of the HER2 molecule comprising at least 9 amino acids, essentially free of transmembrane and intracellular portions of said HER2 molecule.
- 2. An extracellular portion of the HER2 molecule comprising an immune epitope, essentially free of transmembrane and intracellular portions of said HER2 molecule.
- 3. The extracellular portion as defined by claim 1, in substantially pure form.
 - 4. The extracellular portion as defined by claim 1, having a purity of at least about 99%.
- 5. The extracellular portion as defined by claim 1,
 wherein said extracellular portion is antigenic in animals.
 - 6. The extracellular portion as defined by claim 1, further comprising the entire extracellular portion of said HER2 molecule.
- 7. The extracellular portion as defined by claim 1, 20 conjugated with a peptide having immunogenic properties.
 - 8. The extracellular portion as defined by claim 7, wherein said peptide comprises an immune epitope.
 - 9. Isolated DNA encoding the extracellular portion as defined by claim 1, terminating upstream of the portion encoding the transmembrane domain of said HER2 molecule.
 - 10. The isolated DNA as defined by claim 9, terminating at least 1 base pair upstream of the portion encoding the transmembrane domain of said HER2 m lecule.

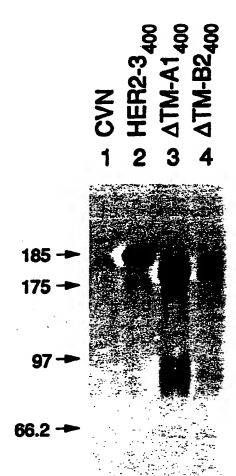
- 11. The isolated DNA as defined by claim 10, terminating about 24 base pairs upstream of the portion encoding the transmembrane domain of said HER2 molecule.
- 12. The isolated DNA as defined by claim 9, wherein said DNA encodes a sequence of at least 9 amino acids of said extracellular portion, and none of the transmembrane or intracellular portions of said HER2 molecule.
 - 13. An expression vector comprising the isolated DNA as defined by claim 9.
- 10 14. The expression vector as defined by claim 13, wherein said expression vector is a virus.
 - 15. A cell into which the expression vector as defined by claim 13 has been introduced.
- 16. The cell as defined by claim 15, wherein said cell is a prokaryote.
 - 17. The cell as defined by claim 15, wherein said cell is a eukaryote.
 - 18. A process for producing an extracellular portion of the HER2 molecule, comprising the steps of:
- a) ligating the isolated DNA as defined by claim 9 into an expression vector capable of expressing said isolated DNA in a suitable host;
 - b) transforming said host with said expression vector;
 - c) culturing said host under conditions suitable for expression of said isolated DNA and production of said extracellular portion; and
 - d) isolating said extracellular porti n fr m said host.

- 19. The process as defin d by claim 18, wherein said host cell is a prokaryote.
- 20. The process as defined by claim 19, wherein said prokaryote is a bacterium.
- 5 21. The process as defined by claim 18, wherein said host cell is a eukaryote.
 - 22. A vaccine comprising the extracellular portion of the HER2 molecule as defined by claim 1.
- 23. The vaccine as defined by claim 22, in combination10 with a suitable adjuvant.
 - 24. A vaccine comprising the extracellular portion of the HER2 molecule as defined by claim 2.
 - 25. The vaccine as defined by claim 24, in combination with a suitable adjuvant.

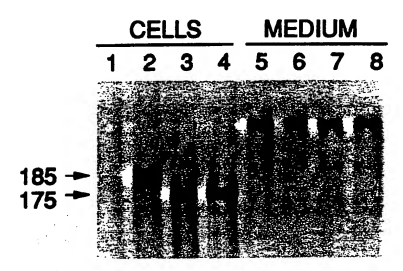




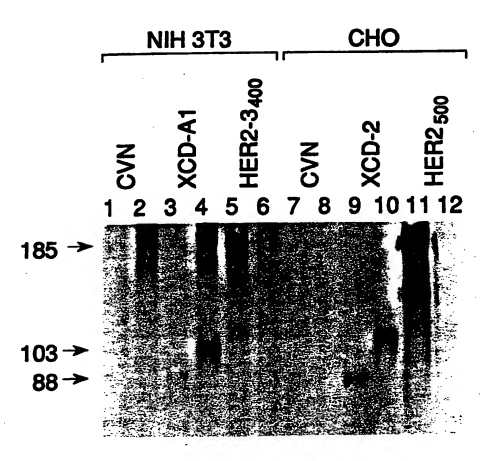
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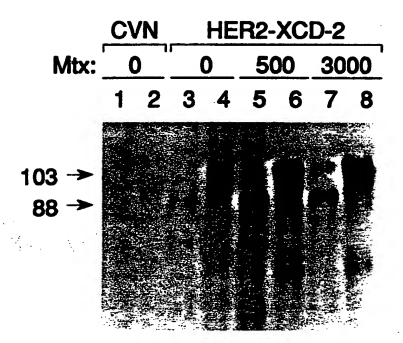
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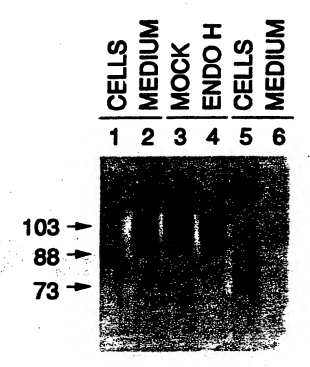
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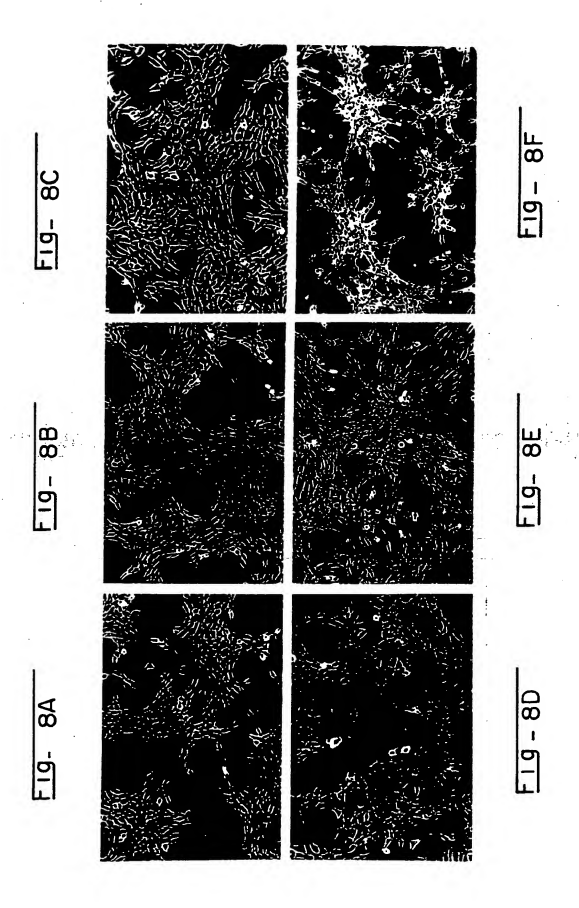
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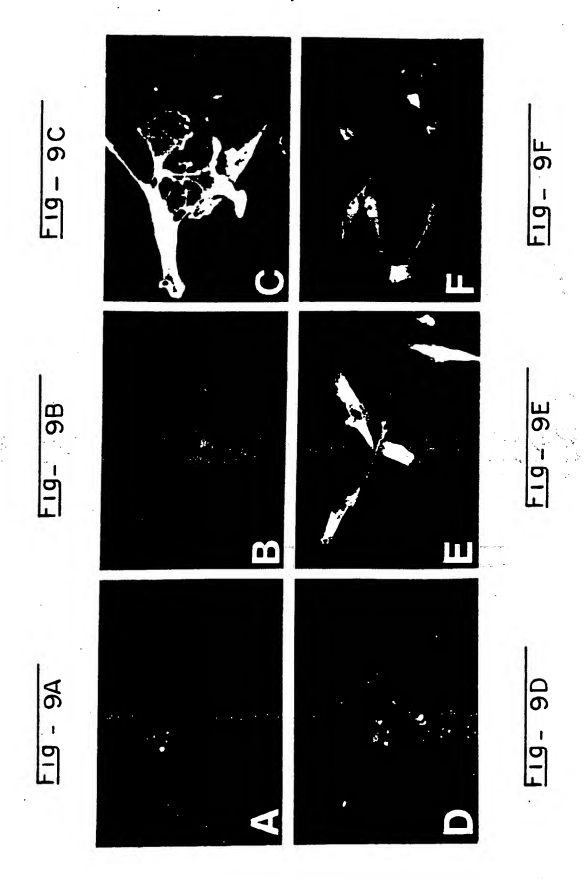
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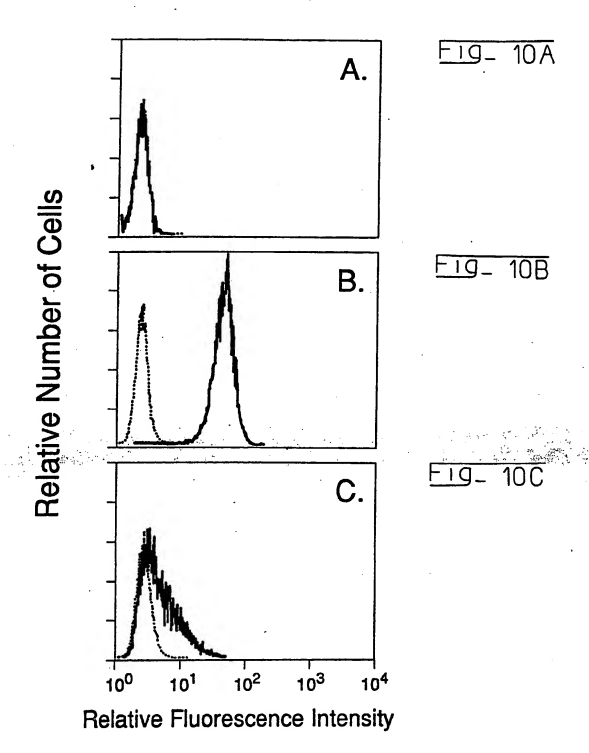
FIG_ 7

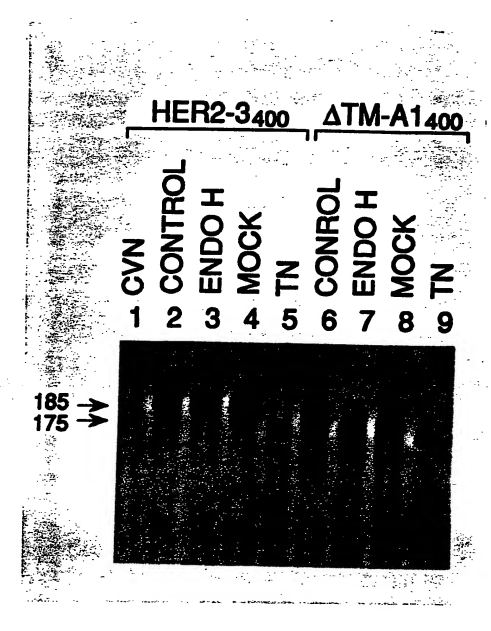


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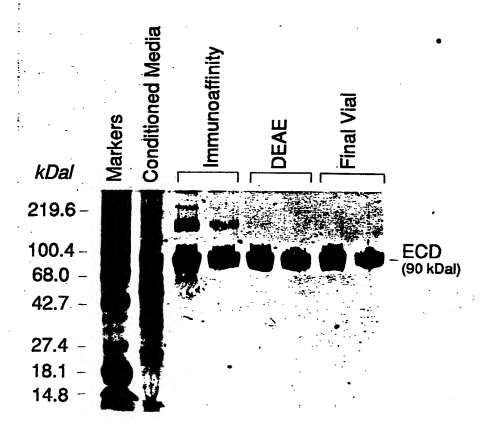




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Purification of the HER2 Extracellular Domain



F19- 12

FIGURE 1

THR	40 GLU GAA	60 GLN CAG	AUG II'S	100 ASP GAC	CBG CBG	140 LEU CUC	160 LEU CUC
GLU	כמפ	VAL	ARG	GLY	COG	GLN	ALA
CCC CCC	ASN	GLU	COC	ASN	GAG	980 000	LEU 1
SER	GEN	GLN	S G	ASP	56 56	ASN 1	GLN 1
ALA	GIN	TLE	CRG	COA	DEC.	ARG 1 CGG 1	ASN C
PRO	VAL	ASP	PRO LEU GLAN ARG LEU CCA CUG CAG AGG CUG	VAL	PRO GLY GLY LEU ARG CCA GGA GGC CUG CGG	GLN 2	ASN A
LEU	VAL	GLN	PRO	ALA GCC	SGA C	ILE (LYS A
ARG	GLN	כתפ	VAL	LEEU COG	PRO	LEU	HIS 1
במפ כמפ	CTS	PHE	GLN	ALA GCC	SER	VAL 1	PHE 1
LYS	GLY CYS	SER	ARG AGG	TYR	ALLA GCC	GGG (ILE I
MET	TYR GLN	50 CUG	70 VAL GUG	90 ASN	110 GLY 2 GGG (130 GLY GGA	150 ASP : GAC :
ASP	TYR	SER	GEN	ASP GAC	ACA	LYS	LYS A
THR	D 20	ALA	ASN	GLU	VAL	בים בים	TRP 1 UGG 1
GGC	LEU ARG HIS LEU CUC CGC CAC CUC	ASN	HIS	PHER	02 D	THR GLU ILE LEU ACA GAG AUC UUG	LEU
ACC	ARG CGC	THR	ALA		JHR ACC	SECO	ILE I
OKS ACKS	D2 22	780 CCC	ILE	GLN	THR	THR CA	THR
VAL	HET	כמפ	בתכ	ACC	ASN	LEU	ASP GAC
S E	ASP	TYR	VAL	GELY	ASN	SER	GLN CAG
ACC ACC	DET	THR	TYR	ARG CGA	COC 1	ARG	TYR C
SER THR AGC ACC	HIS	LEU	299	VAL	PRO	DEC	CYS 1

180 GLY GGC	200 ALA GCC	220 ALA GCU	240 Ser Agu	260 SER UCC	280 TYR UAC	300 GLU GAG	320 Val Gug	340 ILE
LYS	CKS	cys Ugu	HIS	GLU	65.0 CC 53.0	GLN	ARG	
cys	VAL	GLN	ASN	PHE	CYS	ASN	ALA GCC	VAL THR SER ALA ASN GUU ACC AGU GCC AAU
MET	ACC	GLU GAG	PHR	THIR	ALA GCC	HIS	CYS	SER
PRO	SE C	HIS	KIS	ASP	THIR	LEU	PRO	ACC
SER	THR	CYS	LEU	THR	VAL	PRO	LYS	GUO
CKS	LEG	CKS	CYS	ASN	CXS	CYS	SER	SCA
PRO CCC	GLN SER LEU CAG AGC CUG	ASP	ALA	TYR	SER	VAL	CYS	ARG
HIS	S C C	THR	LEU	THR	ALA	COC	LYS	VAL
CYS	CYS	000 000 000	CYS	VAL	399 860	ACC A	35	LEU ARG GLU VAL ARG ALA UUG CGA GAG GUG AGG GCA
NIA GCC	ASP GAU	210 1250 CUG	230 ASP GAC	250 1.60 COG	270 PHE UUC	290 CXS UGC	310 CXS UGU	ARG CGA
ARG CGG	GAG	PRO	SER	ALA	ACA ACA	SER UCC	ARG CGG	LEG
SER	SER	GLY GGG	HIS	PRO	TYR UAU	GEN	CAG	CAS
ARG	SER	LYS	LYS	CYS	ARG CGG	VAL	ACA	GEG
ASN	GLU	CYS	PRO CCC	HIS	GLY	ASP	S S S	MET
THR	GLY	ARG	OCC ALL	LEU	GLU	THR	ASPGAU	SCC
ASP	TRP	ALA	THR	GLU	PRO	SER	GLU	כמפ
ILE	CYS	CYS	CXS	CXS	ASN	Cede	ALA GCA	GGU
LEG	ARG	299 779	OGC GEV	TLE	PRO CCC	TYR	THR ACA	TXR
ACA	SKR	GELY	ALA	GELY	METAUG	ASN	VAL	CYS

FIGURE 1

360 PHE UUU	380 GLU GAG	400 ASP GAC	420 TYR UAC	LEGU CUG	460 PRO CCC	480 GLU GAG	500 TRP UGG	520 VAL GUG
SER	EE	PRO	ALA	OFFI GERN	WAL GUG	PRO CCA	CYS	CYS
GLU	VAL	DE S	GELY	ARG	THR	ARG	HIS	GLU
PRO	GEN	SER	ASN	LIRU	HIS	ASN	GGG	GLN
LEU	COC	ASP	HIS	SKR	VAL	ALA	ARG	GLY
PHE	CAG	PRO	במפ במפ	ARG	PHR	ACO	ALA	ARG
ALA GCA	GLU	TRP	ILE AUU	LEU	CYS	HIS	CYS	כממ
LEU	PRO	AT SO	ARG	555 715	LEU	CAC	LEKU	PHE
SER	GLN	SER	GLY	LEU	HIS	COG	GLN	GLN
999 877	COC	ILE	ARG CGG	TRP	THR	ALA	HIS	SER
PHE UUU	370 PRO CCG	390 TYR UAC	410 ILE AUC	430 SER AGC	450 ASN AAC	470 GLN CAA	490 CYS UGC	510 CYS UGC
RUC	ALA	CUA	VAL	ILE	HIS	HIS	ALA	ASN
EXS	THE	TYR	GLA	GEC	HIS	PRO CCG	LEU	VAL
CYS LYS UGC AAG	ASN	GGU	LEU	LEG	ILE	ASN	GGC	CYS
CYS	SER	THR	ASN	GLY	COC	ARG	GLU	GLN
GEC	ALA	FILE	S G G	GEN	ALA GCC	PHE	299 873	THR
ALA GCU	PRO	GLU ILE GAG AUC	PHR	כנופ	במפ	כמכ	VAL	PRO
PHE	ASP	GEO	VAL	THR	GGA	GLA	CYS	658 666
GLU	GGG	CAG	SER	כתפ	SER	ASP	GLU	PRO
GLN	ASP GAU	ACC	COC	SER	¥15	TRP	ASP	GEU
,,		, = 1	•					

	540 LEU UUG	560 ALA GCU	580 PRO CCC	600 GLY GGC	620 GLY GGC	
	CYS	GLU	CYS	GLU	LYS	
	HIS	PR0 CC6	ARG	GLU	ASP GAC	
	ARG	GEN	ALA GCC	ASP	asp gau	
	ALA SCC	PHEUOU	VAL	PRO CCA	LEG	
	ASIN	CYS	CYS	PHR	ASP	
	VAL	THR	PHE	LYS	VAL	
1	TYR	VAL	PRO	TRP UGG	CYS	
RE 13	GLU	SER	PRO	ILE	SER	
FIGURE	ARG	000 000	ASP	PRO	HIS	
	530 PRO CCC	550 ASN AAU		590 Met Aug	610 THR ACC	**
		GLN	TYR	TYR	CYS	•
	GGG	PRO	HIS	SER	ASN	
	GILN	GLN	ALA	LEU	ILE	
	123	CYS	CYS	ASP	PRO	
	VAL	GLU	ALA	SEC DO	CYS	
	ARG	PRO CCU	VAL ALA GUG GCC	LYS	PRO	624 GLU GAG
	CYS	HIS	CYS	VAL	GLN	ALA
	GLU GLU CYS ARG VAL LEU GLN GLY LEU GAG GAA UGC CGA GUA CUG CAG GGG CUG	CKS	GLAG	gen gen	CYS GLN	CYS PRO ALA UCC CCC GCC
	GLU	PRO	ASP	SER	ALA SCA	CYS

INTERNATIONAL SEARCH REPORT

International Application No

PCT/US90/02697

1. CLASSIFICATION F SUBJECT MATTER (if several classification symbols apply, indicate all) 3						
According to International Patent Classification (IPC) or to both National Classification and IPC						
	5): CO7K 7/06, 7/08, 13/00, 17	/00; Cl2N 15/12, 1/19,	1/21 5/10, 15/63			
C121	21/02; A61K 39/00, 39/385, 39/	38. U.S.Cl. 530/324-328	. 350. 403. 828:			
II. FIELD	S SEARCHED 435/320, 172, 69.1, 6	9.3, 252.3, 240.2; 935/	12. 22-32. 66-75			
		entation Searched 4 424/88				
Classificati		Classification Symbols				
						
	530/324-328, 350, 403, 8		1, 69.3,			
U.S.C	252.3, 240.2; 935/12, 22	-32, 66-75; 424/88				
	Documentation Searched other					
	to the Extent that such Document	s are included in the Fields Searched 5				
Biosi 1975-	s, World Patents Index, USPTO A 1990). See attachment for sear	utomated Patent System ch terms.	(File US PAT,			
III. DOCL	MENTS CONSIDERED TO BE RELEVANT !*		· · · · · · · · · · · · · · · · · · ·			
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Y	Issued 16 January 1986, Yamamot	to et al, "Similarity	3,4,7,8,13-25			
	of Protein Encoded by the Human	n C-erh-B-2 Gene to	1			
	Epiderimal Growth Factor Recept	tor", pages 230-234.				
	See Figures 1-3 and final parag	graph.	:			
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Y	US, A, 4,761,371 (BELL ET AL)	02 August 1988.	7,8,13-21			
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	than the priority date claimed	"A" document member of the same p	atent lamny			
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ATTACHMENT TO PCT/US90/02697

SEARCH TERMS

her2, her 2, Ng1, oncogen?, receptor, erb#, erb b, vaccine, pure, purif?, epidermal growth factor, egf, extra cell?, eternal, ligand, domain, domains, sequence, C erb B 2, erbb2

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